



## CLEAN VERSION OF ALL PENDING CLAIMS AFTER AMENDMENT IS ENTERED

- 1. A process for detecting chromosomal overrepresentation in cells, comprising the following steps:
  - (a) isolating DNAs from cells which have no known numerical changes in their DNAs, and amplifying the DNAs by means of a PCR method using tag primers;
  - (b) hybridizing cells under study in situ with the amplified DNAs from (a);
  - (c) amplifying DNAs from the *in situ* hybridized cells from (b) by means of a PCR method using the tag primers from (a);
  - (d) cohybridizing the DNAs from (a) and (c) to metaphase chromosome spreads from normal cells under suppression hybridization conditions; and
  - (e) identifying numerical changes in the amplified DNAs from (c).
- 2. The process according to claim 1, wherein the cells under study originate from tumors.
- 3. The process according to claim 1, wherein the cells under study originate from the blood of pregnant persons.
- 4. The process according to claim 2 or 3, wherein the cells under study are those of a cell population or single cells.
- 5. The process according to claim 1, wherein the cells under study have an interphase nucleus.

Application Serial No. 09/171,854 Attorney's Docket No. 03528.0038.US00

- 6. The process according to claim 1, wherein the tag primers are degenerative primers.
- 7 The process according to claim 1, wherein the identification from (d) comprises a "Comparative Genomic Hybridization" (CGH) method.
- 8. A kit for carrying out the process according to claim 1, comprising the following components: